L10

L11

89 S L6

E LABRIE F/IN

9 S E4 AND L10

(FILE 'HOME' ENTERED AT 10:47:48 ON 27 JAN 2005) FILE 'CAPLUS' ENTERED AT 10:47:56 ON 27 JAN 2005 2 S WO9626201/PN L1SELECT L1 2 RN L237880 S E1-E51 FILE 'REGISTRY' ENTERED AT 10:49:06 ON 27 JAN 2005 1 S 182167-02-8/RN L3 SET NOTICE 1 DISPLAY SET NOTICE LOGIN DISPLAY FILE 'CAPLUS' ENTERED AT 10:51:44 ON 27 JAN 2005 L4 124379 S ESTROGEN OR ?ESTRADIOL OR ?ESTRIOL OR MESTRANOL L5 70 S L3 OR ACOLBIFENE OR EM(W)652 OR SCH(W)57068 L6 54 S L4(L)L5 L7 24 S L6 NOT PY>=2002 FILE 'MEDLINE, BIOSIS, EMBASE, SCISEARCH' ENTERED AT 11:02:22 ON 27 JAN 2005 L869 S L7 25 DUP REM L8 (44 DUPLICATES REMOVED) L9 FILE 'USPATFULL, USPAT2' ENTERED AT 11:16:22 ON 27 JAN 2005

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=> s wo9626201/pn
             2 WO9626201/PN
=> select l1
ENTER ANSWER NUMBER OR RANGE (1-):2
ENTER DISPLAY CODE (TI) OR ?:rn
E1 THROUGH E51 ASSIGNED
=> s e1-e51
         14732 108-46-3/BI
          3617 110-87-2/BI
          7945 123-08-0/BI
            18 130064-21-0/BI
             9 151533-32-3/BI
            23 151533-34-5/BI
          2032 156-38-7/BI
            52 17720-60-4/BI
          3938 18162-48-6/BI
            49 182167-02-8/BI
            79 182167-03-9/BI
            10 182167-04-0/BI
             3 182167-05-1/BI
             3 182167-06-2/BI
             2 182167-07-3/BI
             2 182167-08-4/BI
             2 182167-09-5/BI
             2 182167-10-8/BI
             2 182167-11-9/BI
             3 182167-12-0/BI
             2 182167-13-1/BI
             2 182167-14-2/BI
             2 182167-15-3/BI
             2 182167-17-5/BI
             2 182167-19-7/BI
             2 182167-21-1/BI
             2 182167-23-3/BI
             2 182167-26-6/BI
             1 182167-28-8/BI
             2 182167-31-3/BI
             2 182167-32-4/BI
             2 182167-34-6/BI
             2 182167-36-8/BI
             2 182167-38-0/BI
             2 182167-39-1/BI
             2 182167-40-4/BI
             2 182167-41-5/BI
             2 182167-43-7/BI
             2 182167-47-1/BI
             9 182167-49-3/BI
             1 182167-53-9/BI
             1 182167-54-0/BI
             1 182167-56-2/BI
             6 182167-58-4/BI
             8 182167-59-5/BI
             2 182330-08-1/BI
           418 1932-03-2/BI
           448 2008-75-5/BI
            43 26815-04-3/BI
          1298 3144-16-9/BI
          4421 3282-30-2/BI
L2
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                OR 18162-48-6/BI OR 182167-02-8/BI OR 182167-03-9/BI OR 182167-
               04-0/BI OR 182167-05-1/BI OR 182167-06-2/BI OR 182167-07-3/BI
               OR 182167-08-4/BI OR 182167-09-5/BI OR 182167-10-8/BI OR 182167-
               11-9/BI OR 182167-12-0/BI OR 182167-13-1/BI OR 182167-14-2/BI
               OR 182167-15-3/BI OR 182167-17-5/BI OR 182167-19-7/BI OR 182167-
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21-1/BI OR 182167-23-3/BI OR 182167-26-6/BI OR 182167-28-8/BI

OR 182167-31-3/BI OR 182167-32-4/BI OR 182167-34-6/BI OR 182167-36-8/BI OR 182167-38-0/BI OR 182167-39-1/BI OR 182167-40-4/BI OR 182167-41-5/BI OR 182167-43-7/BI OR 182167-47-1/BI OR 182167-49-3/BI OR 182167-53-9/BI OR 182167-54-0/BI OR 182167-56-2/BI OR 182167-58-4/BI OR 182167-59-5/BI OR 182330-08-1/BI OR 1932-03-2/BI OR 2008-75-5/BI OR 26815-04-3/BI OR 3144-16-9/BI OR 3282-30-2/BI)

## => SET NOTICE 1 DISPLAY

NOTICE SET TO 1 U.S. DOLLAR FOR DISPLAY COMMAND SET COMMAND COMPLETED

## => D L3 SQIDE 1-

YOU HAVE REQUESTED DATA FROM 1 ANSWERS - CONTINUE? Y/(N):y THE ESTIMATED COST FOR THIS REQUEST IS 6.15 U.S. DOLLARS DO YOU WANT TO CONTINUE WITH THIS REQUEST? (Y)/N:y

- L3 ANSWER 1 OF 1 REGISTRY COPYRIGHT 2005 ACS on STN
- RN 182167-02-8 REGISTRY
- CN 2H-1-Benzopyran-7-ol, 3-(4-hydroxyphenyl)-4-methyl-2-[4-[2-(1-piperidinyl)ethoxy]phenyl]-, (2S)- (9CI) (CA INDEX NAME)

OTHER CA INDEX NAMES:

CN 2H-1-Benzopyran-7-ol, 3-(4-hydroxyphenyl)-4-methyl-2-[4-[2-(1-piperidinyl)ethoxy]phenyl]-, (S)-

OTHER NAMES:

- CN Acolbifene
- CN EM 652
- CN Sch 57068
- FS STEREOSEARCH
- MF C29 H31 N O4
- CI COM
- SR CA
- LC STN Files: ADISINSIGHT, BIOSIS, BIOTECHNO, CA, CAPLUS, EMBASE,
  IMSPATENTS, IMSRESEARCH, PROUSDDR, SYNTHLINE, TOXCENTER, USAN, USPAT2,
  USPATFULL
- DT.CA Caplus document type: Dissertation; Journal; Patent
- RL.P Roles from patents: BIOL (Biological study); PREP (Preparation); RACT (Reactant or reagent); USES (Uses)
- RLD.P Roles for non-specific derivatives from patents: BIOL (Biological study); USES (Uses)
- RL.NP Roles from non-patents: ANST (Analytical study); BIOL (Biological study); MSC (Miscellaneous); PREP (Preparation); PROC (Process); PRP (Properties); RACT (Reactant or reagent); USES (Uses)

Absolute stereochemistry. Rotation (+).

## \*\*PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT\*\*

- 49 REFERENCES IN FILE CA (1907 TO DATE)
- 2 REFERENCES TO NON-SPECIFIC DERIVATIVES IN FILE CA
- 49 REFERENCES IN FILE CAPLUS (1907 TO DATE)

L7 ANSWER 16 OF 24 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 1999:816021 CAPLUS

DOCUMENT NUMBER: 132:117704

TITLE: The anti-estrogen hydroxytamoxifen is a potent

antagonist in a novel yeast system

AUTHOR(S): Liu, Jia Wei; Jeannin, Elisabeth; Picard, Didier

CORPORATE SOURCE: Dep. Biologie Cellulaire, Univ. Geneve, Geneva,

CH-1211, Switz.

SOURCE: Biological Chemistry (1999), 380(11), 1341-1345

CODEN: BICHF3; ISSN: 1431-6730

PUBLISHER: Walter de Gruyter GmbH & Co. KG

DOCUMENT TYPE: Journal LANGUAGE: English

REFERENCE COUNT: 26 THERE ARE 26 CITED REFERENCES AVAILABLE FOR THIS

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L7 ANSWER 17 OF 24 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 1999:580516 CAPLUS

DOCUMENT NUMBER: 131:306858

TITLE: The interaction of raloxifene and the active

metabolite of the antiestrogen EM-800 (SC 5705) with

the human estrogen receptor

AUTHOR(S): Schafer, Jennifer I. MacGregor; Liu, Hong; Tonetti,

Debra A.; Jordan, V. Craiq

CORPORATE SOURCE: Robert H. Lurie Comprehensive Cancer Center,

Northwestern University Medical School, Chicago, IL,

60611, USA

SOURCE: Cancer Research (1999), 59(17), 4308-4313

CODEN: CNREA8; ISSN: 0008-5472

PUBLISHER: AACR Subscription Office

DOCUMENT TYPE: Journal LANGUAGE: English

REFERENCE COUNT: 38 THERE ARE 38 CITED REFERENCES AVAILABLE FOR THIS

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L7 ANSWER 18 OF 24 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 1999:437555 CAPLUS

DOCUMENT NUMBER: 131:208327

TITLE: EM-652 (SCH 57068), a third generation SERM acting as

pure antiestrogen in the mammary gland and endometrium

AUTHOR(S): Labrie, Fernand; Labrie, Claude; Belanger, Alain; Simard, Jacques; Gauthier, Sylvain; Luu-The, Van:

Simard, Jacques; Gauthier, Sylvain; Luu-The, Van; Merand, Yves; Giguere, Vincent; Candas, Bernard; Luo,

Shouqi; Martel, Celine; Singh, Shankar Mohan; Fournier, Marc; Coquet, Agnes; Richard, Virgile; Charbonneau, Ronald; Charpenet, Gilles; Tremblay, Andre; Tremblay, Gilles; Cusan, Lionel; Veilleux,

Raymonde

CORPORATE SOURCE: Oncology and Molecular Endocrinology Research Center,

Centre Hospitalier Universitaire de Quebec (CHUQ),

Pavilion CHUL, Department of Medicine, Laval

University, Quebec, QC, G1V 4G2, Can.

SOURCE: Journal of Steroid Biochemistry and Molecular Biology

(1999), 69(1-6), 51-84

CODEN: JSBBEZ; ISSN: 0960-0760

PUBLISHER: Elsevier Science Ltd.

DOCUMENT TYPE: Journal; General Review

LANGUAGE: English

REFERENCE COUNT: 224 THERE ARE 224 CITED REFERENCES AVAILABLE FOR

THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE

FORMAT

L7 ANSWER 19 OF 24 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 1998:320183 CAPLUS

DOCUMENT NUMBER: 129:62418

TITLE: Binding characteristics of novel nonsteroidal

antiestrogens to the rat uterine estrogen receptors

AUTHOR(S): Martel, Celine; Provencher, Louis; Li, Xun; St. Pierre, Alain; Leblanc, Gilles; Gauthier, Sylvain;

Merand, Yves; Labrie, Fernand

CORPORATE SOURCE: Laboratory of Molecular Endocrinology, CHUL Research

Center, QC, G1V 4G2, Can.

SOURCE: Journal of Steroid Biochemistry and Molecular Biology

(1998), 64(3-4), 199-205

CODEN: JSBBEZ; ISSN: 0960-0760

PUBLISHER: Elsevier Science Ltd.

DOCUMENT TYPE: Journal LANGUAGE: English

REFERENCE COUNT: 33 THERE ARE 33 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L7 ANSWER 20 OF 24 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 1998:153797 CAPLUS

DOCUMENT NUMBER: 128:266369

TITLE: Ligand-independent activation of the estrogen

receptors  $\alpha$  and  $\beta$  by mutations of a

conserved tyrosine can be abolished by antiestrogens AUTHOR(S): Tremblay, Gilles B.; Tremblay, Andre; Labrie, Fernand;

Giguere, Vincent

CORPORATE SOURCE: Molecular Oncology Group, Royal Victoria Hospital,

Montreal, QC, H3A 1A1, Can.

SOURCE: Cancer Research (1998), 58(5), 877-881

CODEN: CNREA8; ISSN: 0008-5472

PUBLISHER: American Association for Cancer Research

DOCUMENT TYPE: Journal LANGUAGE: English

L7 ANSWER 21 OF 24 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 1998:11899 CAPLUS

DOCUMENT NUMBER: 128:149270

TITLE: EM-800, a novel antiestrogen, acts as a pure

antagonist of the transcriptional functions of

estrogen receptors  $\alpha$  and  $\beta$ 

AUTHOR(S): Tremblay, Andre; Tremblay, Gilles B.; Labrie, Claude;

Labrie, Fernand; Giquere, Vincent

CORPORATE SOURCE: Molecular Oncology Group, Royal Victoria Hospital,

Montreal, QC, H3A 1A1, Can.

SOURCE: Endocrinology (1998), 139(1), 111-118

CODEN: ENDOÃO; ISSN: 0013-7227

PUBLISHER: Endocrine Society

DOCUMENT TYPE: Journal LANGUAGE: English

REFERENCE COUNT: 43 THERE ARE 43 CITED REFERENCES AVAILABLE FOR THIS

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L7 ANSWER 22 OF 24 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 1997:693477 CAPLUS

DOCUMENT NUMBER: 128:10399

TITLE: Characterization of the effects of the novel

non-steroidal antiestrogen EM-800 on basal and estrogen-induced proliferation of T-47D, ZR-75-1 and

MCF-7 human breast cancer cells in vitro

AUTHOR(S): Simard, Jacques; Labrie, Claude; Belanger, Alain; Gauthier, Sylvain; Singh, Shankar M.; Merand, Yves;

Talana Daniel, Sylvain, Singh, Shankai M., Melanu

Labrie, Fernand

CORPORATE SOURCE: Laboratory of Molecular Endocrinology, CHUL Research

Center, QC, Can.

SOURCE: International Journal of Cancer (1997), 73(1), 104-112

CODEN: IJCNAW; ISSN: 0020-7136

PUBLISHER: Wiley-Liss DOCUMENT TYPE: Journal

LANGUAGE: English

REFERENCE COUNT: 35 THERE ARE 35 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 23 OF 24 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER:

1997:560295 CAPLUS

DOCUMENT NUMBER:

127:242889

TITLE:

Blockade of the stimulatory effect of estrogens, OH-tamoxifen, OH-toremifene, droloxifene, and raloxifene on alkaline phosphatase activity by the

antiestrogen EM-800 in human endometrial

adenocarcinoma Ishikawa cells

AUTHOR (S):

Simard, Jacques; Sanchez, Rocio; Poirier, Donald; Gauthier, Sylvain; Singh, Shankar M.; Merand, Yves; Belanger, Alain; Labrie, Claude; Labrie, Fernand Laboratory of Molecular Endocrinology, CHUL Research

CORPORATE SOURCE:

Center, Quebec, QC, GIV 4G2, Can.

SOURCE:

Cancer Research (1997), 57(16), 3494-3497

CODEN: CNREA8; ISSN: 0008-5472

PUBLISHER:

American Association for Cancer Research

DOCUMENT TYPE:

Journal English

LANGUAGE: REFERENCE COUNT:

40 THERE ARE 40 CITED REFERENCES AVAILABLE FOR THIS

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 24 OF 24 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER:

1997:296289 CAPLUS

DOCUMENT NUMBER:

127:31462

TITLE:

Response of symbiotic endomycorrhizal fungi to

estrogens and antiestrogens

AUTHOR (S):

Poulin, Marie-Josee; Simard, Jacques; Catford,

Jean-Guy; Labrie, Fernand; Piche, Yves

CORPORATE SOURCE:

Centre de Recherche en Biologie Forestiere, Faculte de

Foresterie et de Geomatique, Universite Laval,

Sainte-Foy, QC, G1K 7P4, Can.

SOURCE:

Molecular Plant-Microbe Interactions (1997), 10(4),

481-487

CODEN: MPMIEL; ISSN: 0894-0282

PUBLISHER:

American Phytopathological Society

DOCUMENT TYPE:

Journal English

LANGUAGE:

REFERENCE COUNT:

46 THERE ARE 46 CITED REFERENCES AVAILABLE FOR THIS

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

was also found to block the recruitment of SRC-1 at AF1 of ERβ, this ligand-independent activation of AF1 being closely. . . protein kinase. Most importantly, the antiestrogen hydroxytamoxifen has no inhibitory effect on the SRC-1-induced ER $\beta$  activity while the pure antiestrogen EM-652 completely abolishes this effect, thus strengthening the need to use pure antiestrogens in breast cancer therapy in order to control. . . up to 5 yr become neg. at longer time intervals and why resistance develops to tamoxifen. EM-800, the prodrug of EM-652, has been shown to prevent the development of dimethylbenz(a)anthracene (DMBA)-induced mammary carcinoma in the rat, a well-recognized model of human. . . Uterine size was reduced to castration levels in the groups of animals treated with EM-800. An almost complete disappearance of estrogen receptors was observed in the uterus, vaginum and tumors in nude mice treated with EM-800. EM -652 was the most potent antiestrogen to inhibit the growth of human breast cancer ZR-75-1, MCF-7 and T-47D cells in vitro when compared with ICI 182780, ICI 164384, hydroxytamoxifen, and droloxifene. Moreover, EM-652 and EM-800 have no stimulatory effect on the basal levels of cell proliferation in the absence of E2 while hydroxytamoxifen and droloxifene had a stimulatory effect on the basal growth of T-47D and ZR-75-1 cells. EM-652 was also the most potent inhibitor of the percentage of cycling cancer cells. When human breast cancer ZR-75-1 xenografts were grown in nude mice, EM-800 led to a complete inhibition of the stimulatory effect of estrogens in ovariectomized mice while tamoxifen was less potent and even stimulated the growth of the tumors in the absence of estrogens, thus illustrating the stimulatory effect of tamoxifen on breast cancer growth. When incubated with human Ishikawa endometrial carcinoma cells, EM-800 had no stimulatory effect on alkaline phosphatase activity, an estrogen -sensitive parameter. Raloxifene, droloxifene, hydroxytoremifene and hydroxytamoxifen, on the other hand, all stimulated to various extent, the activity of this enzyme.. . . tamoxifen failure patients where  ${\tt EM-800}$ (SCH 57050) is compared to Arimidex. The detailed information obtained at the preclin. level with EM-652 or EM-800 indicates that these orally active compds. are highly potent and pure antiestrogens in the mammary gland and endometrium. Estrogens RL: ADV (Adverse effect, including toxicity); BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); THU (Therapeutic use); BIOL (Biological study); USES (Uses) (antiestrogens; activity of EM-652 (SCH 57068) antiestrogen in mammary gland and endometrium) Estrogen receptors RL: BPR (Biological process); BSU (Biological study, unclassified); BIOL (Biological study); PROC (Process)  $(\alpha \text{ and } \beta; \text{ activity of }$ EM-652 ( SCH 57068) antiestrogen in mammary gland and endometrium)

L7 ANSWER 19 OF 24 CAPLUS COPYRIGHT 2005 ACS on STN

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AΒ

Tamoxifen (TAM), the only antiestrogen currently available for the endocrine therapy of breast cancer behaves as a mixed agonist/antagonist of estrogen action, thus limiting its therapeutic potential. We report the binding characteristics of a novel series of nonsteroidal antiestrogens to the rat uterine estrogen receptor. As measured by competition studies, the affinity of EM-652, the active metabolite of the prodrug EM-800, for the estrogen receptor is 7-11 times higher than that of  $17\beta$ - estradiol (E2), ICI 182780, and hydroxy-tamoxifen (OH-TAM), the active metabolite of Tamoxifen. EM-652 is 20+ more potent than ICI 164384 and Droloxifene while it is 400 times more potent than Toremifene in displacing [3H]E2 from the rat uterine estrogen receptor. On the other hand, the prodrug EM-800 and Tamoxifen have resp. 150-fold and 410-fold less affinity for the estrogen receptor than the pure antiestrogen EM-652. No significant binding of EM-652, EM-800, TAM or OH-TAM was observed to the rat uterine progesterone receptor at concns. up to 10 000 nM except for TAM that caused a 50% displacement of labeled R5020 at 4000 nM. No

significant binding of EM-652 or EM-800 was observed on the rat ventral prostate androgen receptor or the rat uterine progesterone receptor. The present data demonstrate the high affinity and specificity of the new antiestrogen, EM-652, for the rat uterine estrogen receptor. The antiestrogen EM-652 thus becomes the compound having the highest known affinity for the estrogen receptor. Due to its unique potency and its pure antiestrogenic activity already demonstrated in many systems, this antiestrogen could well offer an important advance for the endocrine therapy of breast cancer, uterine cancer, and other estrogen -sensitive diseases in women. 50-28-2,  $17\beta$ -Estradiol, biological studies 10540-29-1, Tamoxifen 68047-06-3, Hydroxy-tamoxifen 82413-20-5, Droloxifene 89778-26-7, 98007-99-9, ICI 164384 129453-61-8, ICI 182780 Toremifene 151533-34-5, EM 343 182167-02-8, EM 652 182167-03-9, EM 800 182167-04-0, EM 762. 182167-49-3, EM 776 182167-58-4, EM 651 RL: BPR (Biological process); BSU (Biological study, unclassified); BIOL (Biological study); PROC (Process) (binding characteristics of nonsteroidal antiestrogens to uterine estrogen receptors) ANSWER 20 OF 24 CAPLUS COPYRIGHT 2005 ACS on STN It has recently been suggested that mutation of a conserved tyrosine to asparagine within the ligand-binding domain of the estrogen receptor (ER)  $\alpha$  confers hormone-independent activation and insensitivity to antiestrogens. In view of the recent discovery of  $ER\beta$  and the development of the novel nonsteroidal antiestrogen EM-800 and its active metabolite EM-652, the authors decided to reexamine this issue by introducing a series of mutations at the conserved tyrosine 537 in  $ER\alpha$  and 443 in  $ER\beta$  and measuring their transcriptional activity in the absence and presence of estradiol and the antiestrogens EM-652, ICI 182,780, and 4-hydroxytamoxifen. As demonstrated previously for ERα, the authors observed that substituting a serine or asparagine but 68047-06-3, 4-Hydroxytamoxifen 129453-61-8, ICI-182780 182167-02-8, EM 652 RL: BAC (Biological activity or effector, except adverse); BPR (Biological process); BSU (Biological study, unclassified); BIOL (Biological study); PROC (Process) (ligand-independent activation of the estrogen receptors  $\alpha$  and  $\beta$  by mutations of a conserved tyrosine can be abolished by antiestrogens) ANSWER 21 OF 24 CAPLUS COPYRIGHT 2005 ACS on STN Estrogens act as potent mitogens in a large number of breast cancers, and the use of estrogen receptor (ER) antagonists is, therefore, considered the endocrine therapy of choice in the management of this disease. We describe the mol. properties of EM-652 , the active metabolite of EM-800, a novel nonsteroidal antiestrogen compound, on the transcriptional functions of  $ER\alpha$  and  $ER\beta$ . Using RT-PCR, . . that both receptors should be considered putative targets for antiestrogen action in the breast. In cotransfection assays using a synthetic estrogen-responsive promoter, EM-652 shows no agonistic activity on  $ER\alpha$  and  $ER\beta$  transcriptional function and blocks the estradiol (E2)-mediated activation of both  $ER\alpha$  and  $ER\beta$ . EM-652 is also very effective in abrogating E2-stimulated  $ER\alpha$  and  $ER\beta$ trans-activation of the pS2 promoter in HeLa cells. EM-652 does not alter binding of ER $\alpha$  and ER $\beta$  to DNA. The Ras-mediated induction of  $ER\alpha$  and  $ER\beta$  transcriptional activity

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182167-02-8, EM 652 182167-03-9, EM-800 RL: BAC (Biological activity or effector, except adverse); BSU (Biological

in the presence of E2 is also completely abolished by EM-

activation of  $ER\alpha$  and  $ER\beta$  by the steroid hormone receptor

652. In addition, EM-652 blocks the E2-dependent

coactivator-1 as well as the in vitro.

study, unclassified); THU (Therapeutic use); BIOL (Biological study); USES (Uses)

(EM-800, a novel antiestrogen, acts as a pure antagonist of the transcriptional functions of estrogen receptors  $\alpha$  and  $\beta)$ 

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ANSWER 22 OF 24 CAPLUS COPYRIGHT 2005 ACS on STN Since estrogens play a predominant role in the development and growth of human breast cancer, antiestrogens represent a logical approach to the treatment of this disease. The present study compares the effects of the novel non-steroidal anti-estrogen EM-800 and related compds. with those of a series of anti-estrogens on basal and  $17\beta$ - estradiol (E2)-induced cell proliferation in human breast cancer cell lines. In the absence of added E2, EM-800 and related compds. failed. . . lines. The stimulation of T-47D cell proliferation induced by 0.1 nM E2 was competitively blocked by a simultaneous incubation with EM-652, EM-800, OH-tamoxifen, OH-toremifene, ICI 182780, ICI 164384, droloxifene, tamoxifen and toremifene at apparent Ki values of 0.015, 0.011-0.017, 0.040-0.054, 0.043,. . . and 0.735 nM, approx., 10 nM and >10 nM, resp. Similar data were obtained in ZR-75-1 and/or MCF-7 cells. Moreover, EM-652 was 6-fold more potent than OH-Tamoxifen in inhibiting the proportion of cycling MCF-7 cells. Our data show that EM-800 and EM-652 are the most potent known antiestrogens in human breast cancer cells in vitro and that they are devoid of the estrogenic activity of OH-tamoxifen and droloxifene suggested by stimulation of cell growth in the absence of estrogens in ZR-75-1 and MCF-7 cells. 10540-29-1, Tamoxifen 68047-06-3, 4-Hydroxy-tamoxifen 89778-26-7, Toremifene Droloxifene 98007-99-9, ICI 164384 110503-62-3, 4-Hydroxy toremifene 129453-61-8, ICI 182780 151533-34-5, EM 343 182167-02-8, EM 652 182167-04-0, EM 762 182167-58-4, EM 651 RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); THU (Therapeutic use); BIOL (Biological study); USES (Uses)

(characterization of effects of novel non-steroidal antiestrogens on basal and estrogen-induced proliferation of T-47D, ZR-75-1 and MCF-7 human breast cancer cells in vitro)

L7 ANSWER 23 OF 24 CAPLUS COPYRIGHT 2005 ACS on STN ΑB . . effects of the novel nonsteroidal pure antiestrogen EM-800 and related compds. with those of a series of antiestrogens on the estrogen-sensitive alkaline phosphatase (AP) activity in human endometrial adenocarcinoma Ishikawa cells. Exposure to increasing concns. of up to 1000 nM EM-800 or its active metabolite EM-652 alone failed to affect basal AP activity. In contrast, incubation with 10 nM (Z)-4-OH-tamoxifen, (Z)-4-OH-toremifene, droloxifene, or raloxifene increased the value of this estrogen-sensitive parameter by 3.3-, 3.5-, 2.2-, and 1.6-fold, resp., a stimulatory effect that was completely reversed by simultaneous exposure to 30 nM EM-800. Moreover, the stimulation of AP activity induced by 1 nM  $17\beta$ - estradiol was completely reversed by EM-800, EM-652, or ICI-182780, at the IC50 value of 1.98, 1.01, and 5.64 nM, resp., whereas the partial blockade exerted by (Z)-4-OH-tamoxifen,. . . 41.0, and 3.74 nM, resp. Thus, as assessed by their activity in the human Ishikawa endometrial carcinoma cells, EM-800 and EM-652 are the most potent known antiestrogens in Ishikawa cells, and, most importantly, they are devoid of the estrogenic activity observed. STantiestrogen estrogen alk phosphatase endometrium adenocarcinoma; tamoxifen alk phosphatase endometrium adenocarcinoma; toremifene alk phosphatase endometrium adenocarcinoma; droloxifene alk phosphatase endometrium adenocarcinoma; rodroloxifene alk phosphatase endometrium adenocarcinoma; EM 800 alk phosphatase endometrium adenocarcinoma; EM 652 alk phosphatase endometrium adenocarcinoma

T 68047-06-3, Hydroxytamoxifen 82413-20-5, Droloxifene 84449-90-1, Raloxifene 110503-62-3 129453-61-8, ICI-182780 182167-02-8, EM 652 182167-03-9

RL: ADV (Adverse effect, including toxicity); BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); THU (Therapeutic use); BIOL (Biological study); USES (Uses) (blockade of stimulatory effect of estrogens, hydroxytamoxifen, hydroxytoremifene, droloxifene, and raloxifene on alkaline phosphatase activity by the antiestrogen EM-800 in human endometrial adenocarcinoma Ishikawa cells)

L7 ANSWER 24 OF 24 CAPLUS COPYRIGHT 2005 ACS on STN Plant flavonoids reported previously to act as mol. signals in the AB arbuscular mycorrhizal (AM) symbiosis are known to bind to estrogen receptors and to exert estrogenic effects on mammalian cells. To further investigate the estrogen-like properties of flavonoids the present study examined whether estrogen and antiestrogens have flavonoid-related functions in AM fungi. were performed in a monoaxenic system with the AM fungi Gigaspora. concns. ranging from 0.01 to 10.0  $\mu M$  shows an estimated EC50 value of 3.26 The present results show that  $17\beta$ - estradiol (III) and II exert similar stimulatory effects in G. intraradices. The agonist effect of II was efficiently suppressed by the new antiestrogen EM -652 (IV), which is also consistent with the possible presence of estrogen-binding sites in AM fungi. 68047-06-3, Hydroxy-tamoxifen 131811-54-6, EM-139 182167-02-8, ΙT

EM 652

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); BIOL (Biological study)

(antiestrogen; response of symbiotic endomycorrhizal fungi to estrogens and antiestrogens)

=> d rn str cn 1-5

L1 ANSWER 1 OF 1265 REGISTRY COPYRIGHT 2005 ACS on STN RN 803618-65-7 REGISTRY

Absolute stereochemistry.

\*\*PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT\*\*

CN Estradiol, 3-benzoate 1-piperidineacetate (8CI) (CA INDEX NAME)

L1 ANSWER 2 OF 1265 REGISTRY COPYRIGHT 2005 ACS on STN

RN 786733-95-7 REGISTRY

\*\*\* STRUCTURE DIAGRAM IS NOT AVAILABLE \*\*\*

\*\*\* USE 'SQD' OR 'SQIDE' FORMATS TO DISPLAY SEQUENCE \*.\*\*

\*\*RELATED SEQUENCES AVAILABLE WITH SEQLINK\*\*

CN Protein E2IG5 (human clone DE10316701-SEQID-611 estradiol-induced)
(9CI) (CA INDEX NAME)

OTHER NAMES:

CN 231: PN: DE10316701 PAGE: 1255 claimed sequence

L1 ANSWER 3 OF 1265 REGISTRY COPYRIGHT 2005 ACS on STN

RN 786732-09-0 REGISTRY

\*\*\* STRUCTURE DIAGRAM IS NOT AVAILABLE \*\*\*

\*\*\* USE 'SQD' OR 'SQIDE' FORMATS TO DISPLAY SEQUENCE \*\*\*

CN DNA (human clone DE10316701-SEQID-122 clone E2IG5 estradiol-induced protein cDNA plus flanks) (9CI) (CA INDEX NAME)

OTHER NAMES:

CN 50: PN: DE10316701 PAGE: 874 claimed DNA

L1 ANSWER 4 OF 1265 REGISTRY COPYRIGHT 2005 ACS on STN

RN 736967-60-5 REGISTRY

CN Estra-1,3,5(10)-triene-3,17-diol (17 $\beta$ )-, compd. with methanol (2:1) (9CI) (CA INDEX NAME)

OTHER NAMES:

CN Estradiol compd. with methanol (2:1)

CM 1

Absolute stereochemistry.

L1 ANSWER 5 OF 1265 REGISTRY COPYRIGHT 2005 ACS on STN RN 668276-95-7 REGISTRY

Absolute stereochemistry.

\*\*PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT\*\*

CN Estra-1,3,5(10)-triene-3,17-diol, 11-chloro-, (11 $\beta$ ,17 $\beta$ )- (9CI) (CA INDEX NAME)

OTHER NAMES:

CN  $11\beta$ -Chloroestradiol

```
=> e labrie f/in
                 LABRIE CRAIG B/IN
          12
E2
            2
                 LABRIE DAVID WILLIAM/IN
            0 --> LABRIE F/IN
E3
                LABRIE FERNAND/IN
           86
E4
            1
                 LABRIE JACQUES/IN
E5
                 LABRIE JACQUES J/IN
            2
E6
                  LABRIE JACQUES JOSEPH/IN
            2
E7
              LABRIE JAMES J/IN
LABRIE JAMES R/IN
LABRIE JEAN PIERRE/IN
LABRIE KIMBERLY D/IN
LABRIE MARCEL/IN
E8
            1
            1
E9
            3
E10
            2
E11
E12
            1
=> s e4 and 110
           9 "LABRIE FERNAND"/IN AND L10
=> d ibib 1-9
L11 ANSWER 1 OF 9 USPATFULL on STN
ACCESSION NUMBER:
                      2004:203920 USPATFULL
TITLE:
                      Medical uses of a selective estrogen receptor modulator
                      in combination with sex steroid precursors
INVENTOR(S):
                      Labrie, Fernand, Sainte-foy, CANADA
PATENT ASSIGNEE(S): Endorecherche, Inc. (U.S. individual)
                            NUMBER KIND DATE
                       ______
PATENT INFORMATION: US 2004157812 A1 20040812 APPLICATION INFO.: US 2003-749981 A1 20031230
                                                        (10)
RELATED APPLN. INFO.: Division of Ser. No. US 1999-330799, filed on 11 Jun
                       1999, GRANTED, Pat. No. US 6670346 Continuation-in-part
                       of Ser. No. US 1998-96284, filed on 11 Jun 1998,
                       GRANTED, Pat. No. US 6465445
DOCUMENT TYPE:
                   . Utility
                      APPLICATION
FILE SEGMENT:
LEGAL REPRESENTATIVE: OSTROLENK FABER GERB & SOFFEN, 1180 AVENUE OF THE
                      AMERICAS, NEW YORK, NY, 100368403
NUMBER OF CLAIMS:
EXEMPLARY CLAIM:
NUMBER OF DRAWINGS:
                       17 Drawing Page(s)
LINE COUNT:
                       2192
CAS INDEXING IS AVAILABLE FOR THIS PATENT.
L11 ANSWER 2 OF 9 USPATFULL on STN
ACCESSION NUMBER:
                   2004:72633 USPATFULL
                       Methods of treating and/or suppressing weight gain
TITLE:
INVENTOR(S):
                       Labrie, Fernand, Quebec, CANADA
                       Deshaies, Yves, Quebec, CANADA
                       Richard, Denis, Quebec, CANADA
                       Martel, Celine, Quebec, CANADA
                       Marette, Andre, Quebec, CANADA
PATENT ASSIGNEE(S):
                       Endorecherche, Inc., CANADA (non-U.S. corporation)
                           NUMBER
                                       KIND DATE
                       -----
                                                         (9) OP dai 10
                       US 6710059 B1 20040323
US 2000-610286 20000706
PATENT INFORMATION:
APPLICATION INFO.:
                            NUMBER DATE
PRIORITY INFORMATION: US 1999-142407P 19990706 (60)
DOCUMENT TYPE:
                      Utility
FILE SEGMENT:
                       GRANTED
PRIMARY EXAMINER:
                      Seaman, D. Margaret
LEGAL REPRESENTATIVE: Ostrolenk, Faber, Gerb & Soffen LLP
NUMBER OF CLAIMS:
                       33
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EXEMPLARY CLAIM:

NUMBER OF DRAWINGS: 8 Drawing Figure(s); 8 Drawing Page(s)

LINE COUNT: 2266

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L11 ANSWER 3 OF 9 USPATFULL on STN

2004:45007 USPATFULL ACCESSION NUMBER:

Methods of treating and/or suppressing insulin TITLE:

resistance

INVENTOR (S): Labrie, Fernand, Quebec, CANADA

Deshaies, Yves, Quebec, CANADA Richard, Denis, Quebec, CANADA Martel, Celine, Quebec, CANADA Marette, Andre, Quebec, CANADA

Endorecherche, Inc. (non-U.S. corporation) PATENT ASSIGNEE(S):

> NUMBER KIND DATE

PATENT INFORMATION: US 2004034000 A1 20040219
APPLICATION INFO.: US 2003-387043 A1 20030310 (10)
RELATED APPLN. INFO.: Division of Ser. No. US 2000-610286, filed on 6 Jul
2000, PENDING

NUMBER DATE

PRIORITY INFORMATION: US 1999-142407P 19990706 (60)

PRIORITY INFORMATION: US 1999-142407P 19990706 (60)

DOCUMENT TYPE: FILE SEGMENT: Utility APPLICATION

LEGAL REPRESENTATIVE: OSTROLENK, FABER, GERB & SOFFEN, LLP, 1180 Avenue of

the Americas, New York, NY, 10036-8403

NUMBER OF CLAIMS: EXEMPLARY CLAIM:

NUMBER OF DRAWINGS: 8 Drawing Page(s)

2230 LINE COUNT:

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L11 ANSWER 4 OF 9 USPATFULL on STN

ACCESSION NUMBER: 2003:337283 USPATFULL

TITLE: Medical uses of a selective estrogen receptor modulator

in combination with sex steroid precursors

INVENTOR(S): Labrie, Fernand, Sainte-foy, CANADA

PATENT ASSIGNEE(S): Endorecherche, Inc., CANADA (non-U.S. corporation)

NUMBER KIND DATE \_\_\_\_\_

PATENT INFORMATION: US 6670346 B1 APPLICATION INFO.: US 1999-330799 20031230 19990611 (9)

RELATED APPLN. INFO.: Continuation-in-part of Ser. No. US 1998-96284, filed

on 11 Jun 1998, now patented, Pat. No. US 6465445

Utility GRANTED DOCUMENT TYPE: FILE SEGMENT:

PRIMARY EXAMINER: Criares, Theodore J. ASSISTANT EXAMINER: Kim, Jennifer

LEGAL REPRESENTATIVE: Ostrolenk, Faber, Gerb & Soffen, LLP

NUMBER OF CLAIMS: 15 EXEMPLARY CLAIM:

NUMBER OF DRAWINGS: 17 Drawing Figure(s); 17 Drawing Page(s)

LINE COUNT: 2384

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L11 ANSWER 5 OF 9 USPATFULL on STN

ACCESSION NUMBER: 2003:93647 USPATFULL

TITLE: Selective estrogen receptor modulators in combination

with estrogens

INVENTOR(S): Labrie, Fernand, Sainte-foy, CANADA

PATENT ASSIGNEE(S): Endorecherche, Inc. (non-U.S. corporation)

NUMBER KIND DATE

PATENT INFORMATION: US 2003065008 APPLICATION INFO.: US 2002-143894

A1 20020509 (10) RELATED APPLN. INFO.: Continuation of Ser. No. US 2001-771180, filed on 26

Jan 2001, PENDING

NUMBER DATE

-----

PRIORITY INFORMATION:

US 2000-178601P 20000128 (60)

DOCUMENT TYPE:

Utility

FILE SEGMENT:

APPLICATION

LEGAL REPRESENTATIVE: OSTROLENK FABER GERB & SOFFEN, 1180 AVENUE OF THE

AMERICAS, NEW YORK, NY, 100368403

NUMBER OF CLAIMS:

1

EXEMPLARY CLAIM: NUMBER OF DRAWINGS:

34 Drawing Page(s)

LINE COUNT:

3036

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L11 ANSWER 6 OF 9 USPATFULL on STN

ACCESSION NUMBER:

2003:57944 USPATFULL

TITLE:

Selective estrogen receptor modulators in combination

with estrogens

INVENTOR(S):

Labrie, Fernand, Sainte-foy, CANADA

PATENT ASSIGNEE(S): Endorecherche, Inc. (non-U.S. corporation)

NUMBER KIND DATE \_\_\_\_\_

-----

PATENT INFORMATION: US 2003040510 A1 20030227 APPLICATION INFO.: US 2001-52824 A1 20011107 (10)

wo claim RELATED APPLN. INFO.: Continuation of Ser. No. US 2001-771180, filed on

Jan 2001, PENDING

NUMBER DATE

PRIORITY INFORMATION: US 2000-178601P 20000128 (60)

DOCUMENT TYPE: Utility

FILE SEGMENT:

APPLICATION

LEGAL REPRESENTATIVE: OSTROLENK FABER GERB & SOFFEN, 1180 AVENUE OF THE

AMERICAS, NEW YORK, NY, 100368403

NUMBER OF CLAIMS:

EXEMPLARY CLAIM: NUMBER OF DRAWINGS:

34 Drawing Page(s)

LINE COUNT:

2854

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L11 ANSWER 7 OF 9 USPATFULL on STN

ACCESSION NUMBER:

2002:344449 USPATFULL

TITLE:

Selective estrogen receptor modulators in combination

with estrogens

INVENTOR(S):

Labrie, Fernand, Sainte-foy, CANADA

PATENT ASSIGNEE(S): Endorecherche, Inc. (non-U.S. corporation)

NUMBER KIND DATE \_\_\_\_\_\_ PATENT INFORMATION:

APPLICATION INFO.:

US 2002198179 A1 20021226 US 2001-52803 A1 20011107 (10)

moter application RELATED APPLN. INFO.: Continuation of Ser. No. US 2001-771180, filed on 26

Jan 2001, PENDING

NUMBER DATE

PRIORITY INFORMATION: US 2000-178601P 20000128 (60)

DOCUMENT TYPE:

Utility

FILE SEGMENT:

APPLICATION

LEGAL REPRESENTATIVE: OSTROLENK FABER GERB & SOFFEN, 1180 AVENUE OF THE

AMERICAS, NEW YORK, NY, 100368403

NUMBER OF CLAIMS: EXEMPLARY CLAIM:

43

NUMBER OF DRAWINGS:

34 Drawing Page(s)

LINE COUNT:

3044

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L11 ANSWER 8 OF 9 USPATFULL on STN

ACCESSION NUMBER:

2002:268744 USPATFULL

TITLE:

Medical uses of a selective estrogen receptor modulator

in combination with sex steroid precursors

INVENTOR (S): Labrie, Fernand, Sainte-foy, CANADA

PATENT ASSIGNEE(S):

Endorecherche, Inc., CANADA (non-U.S. corporation)

NUMBER KIND DATE ....., US 6465445 B1 20021015 PATENT INFORMATION: US 1998-96284 APPLICATION INFO.: 19980611 DOCUMENT TYPE: Utility GRANTED FILE SEGMENT:

PRIMARY EXAMINER: Criares, Theodore J. ASSISTANT EXAMINER: Kim, Jennifer

LEGAL REPRESENTATIVE: Ostrolenk, Faber, Gerb & Soffen LLP

NUMBER OF CLAIMS: EXEMPLARY CLAIM:

28

NUMBER OF DRAWINGS:

17 Drawing Figure(s); 13 Drawing Page(s)

LINE COUNT:

2377

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L11 ANSWER 9 OF 9 USPATFULL on STN

ACCESSION NUMBER:

2000:57797 USPATFULL

TITLE:

Benzopyran-containing compounds and method for their

INVENTOR(S):

Labrie, Fernand, Quebec, Canada Merand, Yves, Quebec, Canada Gauthier, Sylvain, Quebec, Canada

PATENT ASSIGNEE(S):

Endorecherche, Inc., Quebec, Canada (non-U.S.

corporation)

NUMBER KIND DATE 20000509 US 6060503

PATENT INFORMATION: APPLICATION INFO.: RELATED APPLN. INFO.:

US 1995-388207 19950221 Continuation-in-part of Ser. No. US 1994-285354, filed

on 3 Aug 1994, now patented, Pat. No. US 5840735 which is a division of Ser. No. US 1991-801704, filed on 2 Dec 1991, now patented, Pat. No. US 5395842

DOCUMENT TYPE:

Utility Granted

FILE SEGMENT: PRIMARY EXAMINER:

Criares, Theodore J.

LEGAL REPRESENTATIVE:

Ostrolenk, Faber, Gerb & Soffen, LLP

NUMBER OF CLAIMS:

EXEMPLARY CLAIM: NUMBER OF DRAWINGS:

5 Drawing Figure(s); 5 Drawing Page(s)

LINE COUNT:

1590

CAS INDEXING IS AVAILABLE FOR THIS PATENT.